## GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: August 3, 2006, 17:48:09; Search time 15911 Seconds

(without alignments)

11490.521 Million cell updates/sec

Title: US-10-761-530-1\_COPY\_442\_3300

Perfect score: 2859

Sequence: 1 atgggagtgaggcacccgcc.....ttctcgtcagctggtgttag 2859

Scoring table: IDENTITY\_NUC

Gapop 10.0 , Gapext 1.0

Searched: 6366136 seqs, 31973710525 residues

Total number of hits satisfying chosen parameters: 12732272

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : GenEmbl:\*

1: gb env:\*

2: gb\_pat:\*

3: gb\_ph:\*

4: gb\_pl:\*

5: gb\_pr:\*

6: gb\_ro:\*

7: gb\_sts:\*

8: gb\_sy:\*

9: gb un:\*

10: gb\_vi:\*

11: gb ov:\*

12: gb\_htg:\*

13: gb\_in:\*

14: gb\_om:\*

15: gb ba:\*

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OM nucleic - nucleic search, using sw model

Run on: August 3, 2006, 17:45:26; Search time 1714 Seconds

(without alignments)

11629.906 Million cell updates/sec

Title: US-10-761-530-1\_COPY\_442\_3300

Perfect score: 2859

Sequence: 1 atgggagtgaggcacccgcc.....ttctcgtcagctggtgttag 2859

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 5244920 seqs, 3486124231 residues

Total number of hits satisfying chosen parameters: 10489840

```
Minimum DB seg length: 0
Maximum DB seq length: 2000000000
Post-processing: Minimum Match 0%
                 Maximum Match 100%
                 Listing first 45 summaries
Database :
                N_Geneseq 8:*
                1: geneseqn1980s:*
                2: geneseqn1990s:*
                3: geneseqn2000s:*
                4: geneseqn2001as:*
                5: geneseqn2001bs:*
                6: geneseqn2002as:*
                7: geneseqn2002bs:*
                8: geneseqn2003as:*
                9: geneseqn2003bs:*
                10: geneseqn2003cs:*
                11: geneseqn2003ds:*
                12: geneseqn2004as:*
                13: geneseqn2004bs:*
                14: geneseqn2005s:*
                15: geneseqn2006s:*
                            GenCore version 5.1.9
                  Copyright (c) 1993 - 2006 Biocceleration Ltd.
OM nucleic - nucleic search, using sw model
                August 3, 2006, 17:56:27; Search time 520 Seconds
Run on:
                                           (without alignments)
                                           10287.500 Million cell updates/sec
Title:
               US-10-761-530-1_COPY_442_3300
Perfect score: 2859
Sequence:
               1 atgggagtgaggcacccgcc.....ttctcgtcagctggtgttag 2859
Scoring table: IDENTITY NUC
               Gapop 10.0 , Gapext 1.0
Searched:
                1403666 seqs, 935554401 residues
Total number of hits satisfying chosen parameters:
                                                       2807332
Minimum DB seq length: 0
Maximum DB seq length: 2000000000
Post-processing: Minimum Match 0%
                Maximum Match 100%
                Listing first 45 summaries
                Issued_Patents_NA:*
Database :
                1: /EMC_Celerra_SIDS3/ptodata/2/ina/1_COMB.seq:*
                2: /EMC_Celerra_SIDS3/ptodata/2/ina/5 COMB.seg:*
                3: /EMC_Celerra_SIDS3/ptodata/2/ina/6A COMB.seq:*
                4: /EMC_Celerra_SIDS3/ptodata/2/ina/6B COMB.seq:*
                5: /EMC_Celerra_SIDS3/ptodata/2/ina/7_COMB.seq:*
                                                                                 6:
/EMC_Celerra_SIDS3/ptodata/2/ina/H_COMB.seq:*
                7: /EMC_Celerra SIDS3/ptodata/2/ina/PCTUS COMB.seg:*
                8: /EMC_Celerra SIDS3/ptodata/2/ina/PP_COMB.seq:*
                9: /EMC_Celerra_SIDS3/ptodata/2/ina/RE COMB.seg:*
                10: /EMC_Celerra_SIDS3/ptodata/2/ina/backfiles1.seg:*
                            GenCore version 5.1.9
```

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OM nucleic - nucleic search, using sw model

Run on: August 3, 2006, 18:04:47; Search time 3002 Seconds

(without alignments)

11702.315 Million cell updates/sec

Title: US-10-761-530-1\_COPY\_442\_3300

Perfect score: 2859

Sequence: 1 atgggagtgaggcacccgcc.....ttctcgtcagctggtgttag 2859

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 18892170 seqs, 6143817638 residues

Total number of hits satisfying chosen parameters: 37784340

Minimum DB seg length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications NA Main:\*

1: /EMC\_Celerra SIDS3/ptodata/2/pubpna/US07 PUBCOMB.seq:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US08\_PUBCOMB.seq:\*

3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US09A PUBCOMB.seq:\*

4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US09B PUBCOMB.seq:\*

i: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US09C\_PUBCOMB.seq:\*

5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10A\_PUBCOMB.seq:\*

7: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10B\_PUBCOMB.seq:\*

8: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10C\_PUBCOMB.seq:\*

9: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10D\_PUBCOMB.seq:\*

10: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10E\_PUBCOMB.seq:\*

11: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10F\_PUBCOMB.seq:\*

12: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10G\_PUBCOMB.seq:\*

13: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US11A\_PUBCOMB.seq:\*

14: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US11B\_PUBCOMB.seq:\*

15: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US11C PUBCOMB.seq:\*

16: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US11D\_PUBCOMB.seq:\*

GenCore version 5.1.9

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OM nucleic - nucleic search, using sw model

Run on: August 4, 2006, 20:23:47; Search time 382 Seconds

(without alignments)

11853.887 Million cell updates/sec

Title: US-10-761-530-1\_COPY\_442\_3300

Perfect score: 2859

Sequence: 1 atgggagtgaggcacccgcc.....ttctcgtcagctqqtqttag 2859

Scoring table: IDENTITY\_NUC

Gapop 10.0 , Gapext 1.0

Searched: 2193277 seqs, 791917567 residues

Total number of hits satisfying chosen parameters: 4386554

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Database : Published\_Applications\_NA\_New:\* 1: /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/US09 NEW PUB.seq:\* 2: /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/US06 NEW PUB.seq:\* 3: /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/US07 NEW PUB.seq:\* 4: /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/US08\_NEW\_PUB.seq:\* /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/PCT\_NEW\_PUB.seq:\* /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/US10\_NEW\_PUB.seq:\* 7: /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/US11\_NEW\_PUB.seq:\* 8: /EMC\_Celerra SIDS3/ptodata/1/pubpna/US11 NEW PUB.seq1:\* 9: /EMC\_Celerra\_SIDS3/ptodata/1/pubpna/US60 NEW PUB.seq:\* GenCore version 5.1.9

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OM nucleic - nucleic search, using sw model

August 3, 2006, 17:54:39; Search time 19980 Seconds Run on:

(without alignments)

8001.670 Million cell updates/sec

Title: US-10-761-530-1\_COPY\_442\_3300

Perfect score: 2859

Sequence: 1 atgggagtgaggcacccgcc.....ttctcgtcagctggtgttag 2859

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 48236798 segs, 27959665780 residues

Total number of hits satisfying chosen parameters: 96473596

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : EST:\*

1: gb\_est1:\* 2: gb\_est3:\* 3: gb\_est4:\* 4: gb\_est5:\* 5: gb\_est6:\* 6: gb\_htc:\* 7: gb\_est2:\* 8: gb\_est7:\* 9: gb\_est8:\* 10: gb\_est9:\*

11: gb\_gss1:\* 12: gb\_gss2:\* 13: gb\_gss3:\* 14: gb\_gss4:\*

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OM protein - protein search, using sw model

August 3, 2006, 13:57:47; Search time 326.791 Seconds Run on:

(without alignments)
1331.951 Million cell updates/sec

Title: US-10-761-530-2

Perfect score: 5105

Sequence: 1 MGVRHPPCSHRLLAVCALVS.....KVLDICVSLLMGEQFLVSWC 952

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2589679 seqs, 457216429 residues

Total number of hits satisfying chosen parameters: 2589679

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : A\_Geneseq\_8:\*

1: geneseqp1980s:\*
2: geneseqp1990s:\*
3: geneseqp2000s:\*
4: geneseqp2001s:\*
5: geneseqp2002s:\*
6: geneseqp2003as:\*
7: geneseqp2003bs:\*

8: geneseqp2004s:\*
9: geneseqp2005s:\*
10: geneseqp2006s:\*

GenCore version 5.1.9
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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:11:03; Search time 71.4245 Seconds

(without alignments)

1166.675 Million cell updates/sec

Title: US-10-761-530-2

Perfect score: 5105

Sequence: 1 MGVRHPPCSHRLLAVCALVS.....KVLDICVSLLMGEQFLVSWC 952

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 650591 seqs, 87530628 residues

Total number of hits satisfying chosen parameters: 650591

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued Patents AA:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/5 COMB.pep:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/6\_COMB.pep:\*

3: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/7\_COMB.pep:\*

4: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/H\_COMB.pep:\*

5: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/PCTUS\_COMB.pep:\*

6: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/RE\_COMB.pep:\*
7: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/backfiles1.pep:\*

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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:27:39; Search time 251.453 Seconds

(without alignments)

1753.728 Million cell updates/sec

Title: US-10-761-530-2

Perfect score: 5105

Sequence:

1 MGVRHPPCSHRLLAVCALVS......KVLDICVSLLMGEQFLVSWC 952

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2097797 seqs, 463214858 residues

Total number of hits satisfying chosen parameters: 2097797

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published\_Applications\_AA\_Main:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US07\_PUBCOMB.pep:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US08\_PUBCOMB.pep:\*

3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US09\_PUBCOMB.pep:\*

4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10A\_PUBCOMB.pep:\*
5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10B\_PUBCOMB.pep:\*

6: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US11\_PUBCOMB.pep:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:28:23; Search time 41.0935 Seconds

(without alignments)

1550.175 Million cell updates/sec

Title: US-10-761-530-2

Perfect score: 5105

Sequence: 1 MGVRHPPCSHRLLAVCALVS.....KVLDICVSLLMGEQFLVSWC 952

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 236815 seqs, 66914042 residues

Total number of hits satisfying chosen parameters: 236815

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database:

Published\_Applications\_AA\_New:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US09\_NEW\_PUB.pep:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US06\_NEW\_PUB.pep:\*

3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US07\_NEW\_PUB.pep:\*

4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US08\_NEW\_PUB.pep:\*

5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/PCT\_NEW\_PUB.pep:\*

6: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10\_NEW\_PUB.pep:\*

7: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US11\_NEW\_PUB.pep:\*

8: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US60\_NEW\_PUB.pep:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:04:39; Search time 50.8777 Seconds

(without alignments)

1800.364 Million cell updates/sec

Title: US-10-761-530-2

Perfect score: 5105

Sequence: 1 MGVRHPPCSHRLLAVCALVS.....KVLDICVSLLMGEQFLVSWC 952

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : PIR\_80:\*

1: pir1:\*
2: pir2:\*
3: pir3:\*
4: pir4:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 13:58:47; Search time 373.755 Seconds

(without alignments)

2356.126 Million cell updates/sec

Title: US-10-761-530-2

Perfect score: 5105

Sequence: 1 MGVRHPPCSHRLLAVCALVS.....KVLDICVSLLMGEQFLVSWC 952

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2849598 seqs, 925015592 residues

Total number of hits satisfying chosen parameters: 2849598

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

UniProt 7.2:\*

1: uniprot\_sprot:\*
2: uniprot\_trembl:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 13:57:47; Search time 7.20863 Seconds

(without alignments)

1331.951 Million cell updates/sec

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Title: US-10-761-530-5

Perfect score: 132

Sequence: 1 MWWRLWWLLLLLLLWPMVWA 21

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2589679 seqs, 457216429 residues

Total number of hits satisfying chosen parameters: 2589679

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : A\_Geneseq\_8:\*

1: geneseqp1980s:\*
2: geneseqp1990s:\*
3: geneseqp2000s:\*
4: geneseqp2001s:\*
5: geneseqp2002s:\*
6: geneseqp2003as:\*
7: geneseqp2003bs:\*
8: geneseqp2004s:\*
9: geneseqp2005s:\*

10: geneseqp2006s:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:11:03; Search time 1.57554 Seconds

(without alignments)

1166.675 Million cell updates/sec

Title: US-10-761-530-5

Perfect score: 132

Sequence:

1 MWWRLWWLLLLLLLWPMVWA 21

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 650591 seqs, 87530628 residues

Total number of hits satisfying chosen parameters: 650591

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued\_Patents\_AA:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/5\_COMB.pep:\*
2: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/6\_COMB.pep:\*
3: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/7\_COMB.pep:\*
4: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/H\_COMB.pep:\*
5: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/PCTUS\_COMB.pep:\*

6: /EMC Celerra\_SIDS3/ptodata/2/iaa/RE COMB.pep:\*

7: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/backfiles1.pep:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:27:39; Search time 5.54676 Seconds

(without alignments)

1753.728 Million cell updates/sec

Title: US-10-761-530-5

Perfect score: 132

Sequence: 1 MWWRLWWLLLLLLLWPMVWA 21

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2097797 seqs, 463214858 residues

Total number of hits satisfying chosen parameters: 2097797

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published\_Applications\_AA\_Main:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US07\_PUBCOMB.pep:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US08\_PUBCOMB.pep:\*

3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US09\_PUBCOMB.pep:\*

4: /EMC\_Celerra SIDS3/ptodata/2/pubpaa/US10A PUBCOMB.pep:\*

5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10B\_PUBCOMB.pep:\*

6: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US11 PUBCOMB.pep:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:28:23 ; Search time 0.906475 Seconds

(without alignments)

1550.175 Million cell updates/sec

Title: US-10-761-530-5

Perfect score: 132

Sequence: 1 MWWRLWWLLLLLLLWPMVWA 21

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 236815 seqs, 66914042 residues

Total number of hits satisfying chosen parameters: 236815

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published\_Applications\_AA\_New:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US09\_NEW\_PUB.pep:\*
2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US06\_NEW\_PUB.pep:\*
3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US07\_NEW\_PUB.pep:\*
4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US08\_NEW\_PUB.pep:\*
5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/PCT\_NEW\_PUB.pep:\*
6: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10\_NEW\_PUB.pep:\*
7: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US11\_NEW\_PUB.pep:\*
8: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US60\_NEW\_PUB.pep:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 14:04:39; Search time 1.1223 Seconds

(without alignments)

1800.364 Million cell updates/sec

Title: US-10-761-530-5

Perfect score: 132

Sequence: 1 MWWRLWWLLLLLLLWPMVWA 21

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : PIR\_80:\*

1: pir1:\* 2: pir2:\* 3: pir3:\* 4: pir4:\*

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: August 3, 2006, 13:58:47; Search time 8.2446 Seconds

(without alignments)
2356.126 Million cell updates/sec

Title: US-10-761-530-5

Perfect score: 132

. . .

Sequence: 1 MWWRLWWLLLLLLLWPMVWA 21

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2849598 seqs, 925015592 residues

Total number of hits satisfying chosen parameters: 2849598

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : UniProt\_7.2:\*

1: uniprot\_sprot:\*
2: uniprot\_trembl:\*